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- (54) VINYL-CIS-POLYBUTADIENE RUBBER AND BUTADIENE RUBBER COMPOSITION USING SAME
- (57) The present invention relates to a viny-cispolybutadiene rubber containing 1,2-polybutadiene and a polymer substance of a melting point lower than that of the 1,2-polybutadiene and with at least one unsaturated double bond per repeating unit, where the 1,2-polybutadiene and the polymer substance are dispersed at physically and/or chemically adorbed states in the cis-

polybutadiene rubber as the matrix component of the vinyl-cis-polybutadiene rubber, and a method for producing the same, to provide a vulcenized product exerting a small die swell ratio and excellent extrusion processability and operability, as well as very great characteristics demanded for the side tread of tire and the like.

Description

Technical Field

5 00011 The present invention relates to a novel virylicia-polybutadeine nubber produced by concurrently allowing 1,2-ophytustadeine of a high melting point of 170°C or more and polybergone or polybutadeine of a low melting point to axist and be dispersed in the matrix of cis-polybutadeine nubber. Further, the invention relates to a butadeine nubber composition using the virtue's colvibutadeine nubber.

10 Background of the Invention

[0002] In the molecular chain of polybutadiane, a binding portion generated by 1.4 polymerization (1.4 structure) and a binding portion generated by 1.4 polymerization (1.4 structure) and to a binding portion generated by 1.2 polymerization (1.2 structure) concurrently exist as so called inforestructure). Attendatively, the 1.2 structure is divided in two types of structures, namely cis structure and trans structure. Alternatively, the 1.2 structure lakes a structure with vinity group as a side chain.

[0003] A method for producing vinyl-cie-polybutadiene rubber composition in the related art has been carried out in inert organic solvents such as aromatic hydrocarbons such as benzene, follower and halogenated hydrocarbons thereof for example chiorobenzene. When solvents such as aromatic hydrocarbons and halogenated hydrocarbons are used, however, the resulting polymerization solution has such a high viscosity that the agitation, heat transmission and transfer thereof are troublesome, which requires excessive energy for the recovery of such solvent. Additionally, solvents such as aromatic hydrocarbons and halogenated hydrocarbons are very hazardous for environment, due to the toxicity and carbinogenesis.

[2004] Ås the production method, a method including a step of producing cis-polybutadene rubber by cis-1, 4 polymerization of 1,3-butadiene using a catalyst obtained from water, a soluble cobatic corpound and an organic autimation of cinidde represented by the general formula Alfi₂X₂₋₈, (provided that R is an alkyl group with one to 6 carbon atoms, phenyl group or cycloskyl group; X is a halogen element; and n is a numerical figure of 1.5 to 2) in the inert organic solvent, and a step of syndicatic 1,2 polymerization (abbrevised as 1² polymerization hereinater) of 1,3-butadene in the presence of a syndicatic 1,2 polymerization catalyst obtained from a soluble cobat compound, an organic autiminum compound represented by the general formula AIFs_(provided that R is an alkyl group with one to 6 carbon of sound, phenyl group or cycloskyl group) and carbon disultide, with addition or no addition of 1,3-butadene and/or the

atoms, phenyl group or cyclosidly group) and carbon disulfide, with addition or no addition of 1,3-butadien and/or the solvent to the resulting polymerization system is known (see for example JP-B-49-17666 (patent reference 1) and JP-B-49-17667 (patent reference 2)).

[0005] Additionally, for example, JP-B-62-171 (patent reference 3), JP-B-63-36324 (patent reference 4), JP-B-2-37927

(patient reference \$), JP-B2-28081 (patient reference \$), and JP-B2-863666 (patient reference \$7 / describe methods including a step of producing vinyl-cis-polybutadiene nebber composition by cis-1,4 polymerization of 1,3-butadiene in the presence or absence of carbon disulfide, and methods including a step of separating and recovering 1,3-butadiene and carbon disulfide in creycle 1,3-butadiene substantially never containing carbon disulfide and the inert organic solvent. Further, JP-34-48816 (patient reference 8) describes a rubber composition with a small dis swall ratio of compounded material, of which the vulcanized product has an excellent tensile strength and a great flex-crack-growth resistance preferable as the sidewall of its and the sidewall of t

[0005] Additionally, JP-A-2000-44833 (patent reference 9) describes a method for producing viryl-cis-polybutadiene rubber in an Inent organic solvent containing C, delitables such as n-butane, cis-2-butane, runs-2-butane, and butene-1 as the main components. JP-A-2000-44833 describes that 1,2-polybutadiene contained in the rubber composition according to the method is a crystal in short fiber, where 98 % or more of the fiber length is less than 0.5 µm in the distribution of the crystal in short fiber along major axis; and 70 % or more thereof is less than 0.2 µm and that the resulting nubber composition has improved moldability of cis-1,4-polybutadiene, tenales letress, tenales strength and flox-

crack-growth resistance.

[0007] However, a rubber composition with improved various properties has been demanded for some use.

Patent Reference 1: JP-B-49-17666
Patent Reference 2: JP-B-49-17667

Patent Reference 3: .IP-R-62-171

Patent Reference 4: .IP-R-63-36324

Patent Reference 5: JP-B-2-37927

Patent Reference 6: JP-B-2-38081

Patent Reference 7: JP-B-3-63566

Patent Reference 8: JP-B-4-48815

Patent Reference 9: JP-A-2000-44633

Disclosure of the Invention

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- [0008] It is an object of the invention to provide vinyl-cles polybutardiene rubber giving a butacdiene rubber composition with a small die sewel ratio and excellent extrusion processability and producing fine, which excets excellent break-resistant properties, abrasion resistance and sliding infection resistance as well as very great fine-crack-growth resistance and high rigidity, when the butacliene rubber composition is vulcanized. It is an object of the invention to provide a butacliene rubber composition with the excellent properties, particularly a butacliene rubber composition for tire. [0009] The invention has achieved the objects via the following constitutions.
- 1. A vinyl cis-polybutadiene rubber containing 1,2-polybutadiene and a polymer substance with a melting point lower than that of the 1,2-polybutadiene and with at least one unsaturated double bond per repeating unit, where the 1,2polybutadiene and the polymer substance are dispersed at physically and/or chemically adsorbed states in the cispolybutadiene rubber as the matrix component of the vinyl-cis-polybutadiene rubber.
 - The vinyt-cis-polybutadiene rubber described in 1. above, where the 1,2-polybutadiene and the polymer substance are dispersed in short crystal fiber and/or particle in the cis-polybutadiene rubber as the matrix component of the vinyt-cis-polybutadiene rubber.
 - 3. The vinty-dis-polybutadiene rubber described in 1. or 2. above, where the 1,2-polybutadiene is 1,2-polybutadiene of a melting-point of 170°C or more and the polymer substance is at least one selected from polybeoprene, crystallizable polybutadiene of a melting point of 150°C or less, liquid polybutadiene and derivatives thereof.
- 4. The vinyl-cis-polybutadiene rubber described in any of 1, through 3, above, where the unsaturated polymer substance is contained within a range of 0.01 to 50 % by mass to the total of the crystal fiber of the 1,2-polybutadiene and cis-polybutadiene rubber.
 - The vinyl-cis-polybutadiene rubber described in any of 1. through 4., where the viscosity of the cis-polybutadiene rubber as the matrix component in toluene solution at 25°C is within a range of 10 to 150.
- The vinyl-cis-polybutadiene rubber described in any of 1. through 5., where [n] of the cis-polybutadiene rubber
 as the matrix component is within a range of 1.0 to 5.0.
 - 7. The vinyl-cis-polybutadiene rubber described in any of 1. through 6., where the content of the 1,4-cis structure of the cis-polybutadiene rubber as the matrix component is within a range of 80 % by mass or more.
 - 8. The vinyl-cls-polybutadiene rubber described in any of 1. through 7, above, where the Mooney viscosity of the cls-polybutadiene rubber as the matrix component of the vinyl-cls-polybutadiene rubber is within a range of 10 to 50.
 9. The vinyl-cls-polybutadiene rubber described in any of 1, through 8, where the polymer substance is a matter insoluble in boiling n-hexane.
 - 10. The vinyl-dis-polybutadiene rubber described in any of 1. Inrough 9., where the 1,2-polybutadiene is dispersable in both crystal fiber in the cis-polybutadiene rubber as the matrix component of the vinyl-dis-polybutadiene rubber as the matrix component of the vinyl-dis-polybutadiene rubber and the polymer substance is dispersed in particle therein, and where the short crystal fiber of the 1,2-polybutadiene is dispersed in the particle of the polymer substance.
 - 11. The viny-rois polybutadiene nubber described in 11 above, where the short crystal fiber of the 1,2-polybutadiene is never contained in the particle of the polymer substance but is also dispersed in the clas-polybutadiene rubber as the matrix component, and where the length of the short crystal fiber dispersed in the matrix along major axis is within a range of 0.2 to 1,000 µm and the length of the short crystal fiber of the 1,2-polybutadiene dispersed in the particle of the polymer substance along major axis is within a range of 0.0 to 10.5 line.
 - 12. A butadiene rubber composition prepared by compounding the vinyl-cis-polybutadiene rubber described in 1. or 2. above at 10 to 300 parts by weight per 100 parts by weight of a rubber selected from natural rubber, polylisoprene rubber, styrene-butadiene copolymer rubber, or a blend rubber of at least two types thereof.
- 49 13. A butadiene rubber composition for tire, where the vinyl-cis-polybutadiene rubber described in 1. through 11. above, and/or the butadiene rubber composition described in 13. above is used. 14. A method for producing vinyl-dis-polybutadiene rubber by a step of the cis-1,4 polymer/zation of 1,3-butadiene
 - using a cis-1,4 polymerization catalysis in a hydrocarbon-series solvent, a step of the 1.2 polymerization of 1,3butadiene in the concurrent presence of a 1,5 polymerization catalyst in the resulting polymerization mixture to generate 1.2-polybutadiene of a melting point of 170°C ormore, and a step of the separation and recovery of virty-toispolybutadiene rubber generated form the resulting polymerization mixture, the method including as tep of adding a polymer substance with at least one unsaturated double bond per repeating unit to the production system of virty-toispolybutadiene rubber.
 - 15. The method for producing vinyl-cis-polybutadiene rubber as described in 14. above, where the polymer substance is at least one selected from polyisoprene, crystallizable polybutadiene of a melting point of 0°C to 150°C, liquid polybutadiene, and derivatives thereof.
 - 16. The method for producing vinyl-cis-polybutadiene rubber as described in 14. or 15. above, where the amount of the polymer substance to be added to the production system is within a range of 0.01 to 50 % by mass to the

vinvi-cis-polybutadiene rubber to be obtained.

- 17. The method for producing vinyl-dis-polybutadione rubber as described in any of 14, through 16, above, where the step of adding the polymer substance to the production system is carried out in the polymerization mixture at an appropriate time point from the step of the cis-14 polymerization step to the step of the separation and recovery of the vinyl-dis-polybutadione rubber generated from the polymerization mixture obtained after the completion of the 12 polymerization.
- 18. The method for producing vinyl-cis-polybutadiene rubber as described in any of 14. through 17. above, where the hydrocarbon-series solvent is a hydrocarbon-series solvent with a solubility parameter of 9.0 or less.
- 19. A bitadiene rubber composition prepared by compounding the vinyl-dis-polybutadiene rubber obtained by the production method described in any of 14. through 18. above at 10 to 300 parts by mass of 20 parts by mass of 20 parts by mass of 20 parts of 20
 - 20. A butadiene rubber composition for tire, where the vinyl-cis-polybutadiene rubber obtained by the production method described in any of 14, through 18, above and/or the butadiene rubber composition described in 12, 13 or 18 above is used.

[0010] In a preferable embodiment, the virty-cis-polydutadiene nubber of the invention (abbreviated as YCRP hereinafter) is a novel YCR where the 1.2-polydutadiene is 1.2-polydutadiene of a melting point of 170°C or more, where the polymer substance of a melting point lower than that of the 1.2-polydutadiene and with at least one unsaturated double bond per repeating unit (sometimes abbreviated as "unsaturated polymer substance" hereinafter) is at least one selected from polysicoprene, organizable polydutadiene of a melting point is est than 170°C, fliquid polydutadiene and derivatives thereor, and where the 1.2-polydutadiene of a melting point of 170°C or more and the unsaturated polymer substance are present concurrently in the matrix of ics-polydutadiene unbaber and are discepted therein.

- [011] Owing to the concurrent presence of 1,2-polybutadiene of a high melting point exerting very strong interaction between polymers as a very excellent reinforcing component and such unsaturated polymer substance with a relatively low melting point such as polysogence, consequently, the VCR in accordance with the invention has remarkably improved dispersibility of the 1,2-polybutadiene of a high melting point in the cis-polybutadiene rubber as the matrix component due to the compatible effect of the concurrently present unsaturated polymer substance, compared with the VCR in the related art, so that the content of the 1,2-polybutadiene of a high melting point as an excellent reinforcing component
- can be raised.
 (Dot2) The characteristics of the VCR in accordance with the invention as described above enable great improvement of various physico-chemical properties strongly demanded in the production of the articles and in other uses. When the VCR of the invention is used in a butadeien rubber composition for the, in particular, the composition has such a small die swell ratio (as the ratio of the dameter of the compounded material to die orifice diameter during extrusion) quiring it improduction, so that the composition exerts great extrusion processability and operability. Additionally, the vulcanized product of the composition exerts excellent break-resistant properties, abrasion resistance, sliding friction resistance and the like mainly required for side tread of the and the like. Because the flex-crack-provid resistance thereof is very great and the rigidity thereof is high, further, the amount of reinforcing materials such as carbon and silica to be used on an be reduced, enabling low fuel consumption owing to the weight decrease of the. Thus, the using the VCR of the invention as a raw material for side treads and the like exerts excellent running stability and high-speed durability and additionally exables low fuel consumption.

Brief Description of the Drawings

45 [0013]

- Fig.1 is one schematic view of a dispersion embodiment of the unsaturated polymer substance in relation to the crystal fiber of the 1,2-polybutadiene of a melting point of 170°C or more.
- Fig.2 is another schematic view of a dispersion embodiment of the unsaturated polymer substance in relation to the crystal fiber of the 1,2-polybutadiene of a melting point of 170°C or more.
 - Fig.3 is a still other schematic view of a dispersion embodiment of the unsaturated polymer substance in relation to the crystal fiber of the 1,2-polybutadiene of a melting point of 170°C or more.
 - Fig.4 is an additional schematic view of a dispersion embodiment of the unsaturated polymer substance in relation to the crystal fiber of the 1,2-polybutadiene of a melting point of 170°C or more.
- Fig.5 is an electron micrograph depicting the microstructure of the vinyl-cis-polybutadiene rubber obtained in Comparative Example 1.
 - Fig. 6 is an electron micrograph depicting the microstructure of the vinyl-cis-polybutadiene rubber obtained in Example

Fig.7 is an electron micrograph depicting the microstructure of the vinyl-cis-polybutadiene rubber obtained in Example

Fig.8 is an electron micrograph depicting the microstructure of the vinyl-cis-polybutadiene rubber obtained in Example

[0014] In the figures, symbol "1" expresses matrix; "2", the crystal fiber of the 1,2-polybutadiene of a melting point of 170°C or more; "3", the microparticle of unsaturated polymer substance.

Best Mode for Carrying out the Invention

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[0015] The VCR of the invention generally has the following constitution. Specifically, the VCR generally contains (1) one to 50 parts by mass of 1,2-polybutadiene of a melting point of 170°C or more; (2) 100 parts by mass of dispolybutadiene rubber and (3) an unsaturated polymer substance at 0.01 to 50 % by mass of the total of (1) and (2). Additionally, the 1,2-polybutadiene of a melting point of 170°C or more as the component (1) generally forms a crystal fiber with the mean length of the mono-dispersed fiber crystal along short axis being 0.2 un or less and an aspect ratio being 10 or less, which is in a short fiber form with the mean number of the mono-dispersed fiber crystal being 10 or more, to the mono-dispersed fiber crystal along short axis being 0.2 µm or less, preferably 0.1 µm or less; with an aspect ratio of 10 or less, preferably 3 or less; and with the mean number of the mono-dispersed fiber crystal along short axis being 0.2 µm or less, preferably 0.1 µm or less; with an aspect ratio of 10 or less, preferably 3 or less; and with the mean number of the mono-dispersed fiber crystal being 10 or more, preferably 15 to 0.220°C.

[0017] The cis-polybutadiene rubber as the component (2) preferably has the following characteristics. In other words, the cis-polybutadiene rubber as the component (2) has a Mooney viscosity (ML_++1 00°C abbreviated as "ML" hereinative of preferably 10 to 40. In such manner, effectively, the operability during compounding is improved, while the dispersion of the component (1) in the component (2) his improved. Additionally, the cis-polybutadiene rubber as the component (2) has preferably the 100 his preferably 10 to 150, more preferably 10 to 100; and [1] (ritrinals viscosity therein to the under the component (2) has preferably 10 to 40. Additionally, the content ratio of the 1.4-bis structure is 80 % by mass or more, preferably 90 % by mass or more. Additionally, the content ratio of the 1.4-bis structure is 80% by the substantially never containing gel matters' means that toluene-insoluble matters are the 30 % by mass or the phrase "substantially never containing gel matters" means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing gel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters' means that toluene-insoluble matters are 1.6 % by mass or the substantially never containing sel matters.

[0018] The end and/or main chain of the polybutadiene rubber obtained by the cls-1,4 polymerization may be modified. As such modifier, organic silicone compounds containing at least amino group and alkoxy group, organic silicone compounds containing alkoxy group, unsaturated carboxylic acid or derivatives thereof, halogen-series compounds, and compounds with hetero-three membered-rings may be used. The amount of such modifier to be used is 0.01 to 150 mmol per 100g of the generated polybutadiene (polybutadiene rubber). When the amount of the modifier to be used is less, the modification affect is hardly exerted. When the amount thereof to be used is too much, the modifier still unreactive is likely to remain in polybutadiene. It requires laborious works to eliminate the modifier, unpreferably. Herein, the Mooney viscosity of the modified product is preferably increased by one or more, compared with the original rubber before modification. So as to promote the reaction, organic peroxides can be added. Preferably, the modified polybutadiene obtained by the method has a Mooney viscosity (ML1+4, 100°C) within a range of 20 to 80 and has a weight average molecular weight of 200,000 to 1,000,000 by gel permeation method, where 80 % by mass or more of the repeating unit has cis-1,4 structure. Additionally, the content of the vinyl structure in the microstructure is preferably 15 % by mass or less. [0019] Herein, the toluene-insoluble matters express gel matters attached on a metal net after filtration, by completely dissolving 10 g of a sample rubber and 400 ml of toluene in an Erlenmeyer flask at RT (25°C) and filtering the resulting solution, using a filtration device arranged with the metal net of 200 mesh. The ratio expresses a value measured by drying the net attached with the gel in vacuum to measure the attached amount thereof to calculate the percentage to the sample rubber.

[0020] Additionally, fill (intrinsic viscosity) is a value determined according to the following formula, by placing o.1 g of a sample rubber and 100 ml of blosen is an Efermeyer flask, completely dissolving the sample rubber at 30°C, and sequently placing the solution of 10 ml in a Canon Fenske dynamic viscometer in a thermostat water bath controlled at 30°C, and measuring the dropt time (f) of the solution.

 $\eta sp = T/T_0 - 1$

(Tn: drop time of toluene alone)

$\eta sp/c = [\eta] + k' [\eta]^2C$

- (ηsp: specific viscosity; k': Huggins constant (0.37); C: sample concentration (g/ml))
- [0021] The rallo between the 1,2-polybutadiene crystal fiber as the component (1) and the cis-polybutadiene as the component (2) is one to 50 parts by mass, preferably one so 30 parts by mass of the 1,2-polybutadiene so of the 1,2-polybutadiene crystal fiber as the component (1) to 100 parts by mass of the cis-polybutadiene as the component (2). Whith the range, the following drawbacks can be avoided: when the amount of the 1,2-polyutadiene crystal fiber is so large to exceed 50 parts by mass, the stort fiber crystal of the 12-polybutadiene crystal fiber in the cis-polybutadiene crystal fiber is small test shan one part by mass, the reinforcibility with the short fiber crystal is deteriorated. Thus, problems hardly occur, such that the characteristic elastic modulus, fixe-crack-cryother hesistance, and oxidation degradation are exerted with difficulty and the processability
- is deteriorated. Therefore, the range is preferable. Further, the ratio of the unsaturated polymer substance as the component (3) is 0.0 to 50% by mass, preferably 0.0 to 50% by mass of VCR, as described above. The range is preferable because the deterioration of the dispersibility due to the aggregation of the 1,2-polytucialene crystal fibers as the component (1) can be suppressed, and an associated deterioration of the various physico-chemical properties of VCR can be suppressed.
- [0022] Further, the ratio of the 1,2-polybutadiene of a melting point of 170°C or more as the component (1) and the unsaturated polymer substance as the component (3) is 0.02 to 100 parts by mass, preferably 0.05 to 80 parts by mass of the component (1) per 100 parts by mass of the component (1). Additionally, the total amount of the components (1) and (3) is 1.01 to 100 parts by mass, preferably 1.03 to 90 parts by mass per 100 parts by mass of the de-polybutadiene nubber as the component (2).
 - [0023] The method for producing VCR in accordance with the invention is described below in detail.
- 5 [00.24] For the VCR production in accordance with the invention, generally, 1.3-butacliene is polymerized, using a hydrocarbon-series selvent. In hydrocarbon-series selvent is preferably a hydrocarbon-series solvent with a solubility parameter (abbreviated as "SP value" hereinater) of 9.0 or less and is more preferably a hydrocarbon-series solvent with a solubility parameter of 9.6 or less includes for example aliphate hydrocarbons and alloycle hydrocarbons, such as hexame (SP value: 7.2), n-pentance (SP value:
- for example aliphatic hydrocarbons and alicyclic hydrocarbons, such as n-hexane (SP value 7.2), n-pentane (SP value 7.0), n-octane (SP value 7.5), cyclohexane (SP value 8.1) and n-butane (SP value 6.6). Among them, for example, cyclohexane is preferable.
 - (0025) The SP values of these solvents are known in references such as Rubber Industry Manual (Gomu Kogyo Blirran) (the 4th edition, Nippon Rubber Association Foundation (Nippon Gomu Kyoka), issued on January 20, 1994, page 721).
- 15 [0026] By using a solvent with an SP value smaller than 9.0, preferably, the dispersion of the short fiber crystal of the 1.2-polybutadiene crystal fiber in the dis-polybutadiene nubber is at a state expected in accordance with the invention, so that excellent die swell characteristic, high tensile stress, tensile strength and high flex-crack-growth performance can be preferably exerted.
- [0027] First, 1,3-butadiene and the solvent are mixed together, to adjust the concentration of water in the resulting solution. Where is within a range of preferably 0 to 1,0 mole particularly preferably 0.2 to 1,0 mole per one mote of an organic aluminum chloride used as the cis-1,4 polymerization catalyst in the solution. The range is preferable because sufficient catalytic activity can be obtained to provide a preferable content and molecular weight of cis-1,4 structure and because get locorurence during polymerization can be suppressed, to prevent glei adhesion note polymerization tanks or the like, so that continuous polymerization time can be protonged. As the method for adjusting water concentration, from the like of the continuous polymerization time can be protonged. As the method of adjusting water concentration, from the like of the continuous polymerization time can be protonged.
- [0028] To the solution obtained by adjusting water concentration is added an organic alumfuum chloride as one of cs. 1.4 polymerization catalysts. As such organic aluminum chloride, a compound represented by the general formula ARI_{N,So.} is preferably used. Specific examples thereof preferably include distriptialuminum monochronide, discipulatiminum discipulatim
 - [0029] Then, a soluble cobalt compound as another one of the cis-1.4 polymerization catalysts is added to a mixture solution to which the organic aluminum chloride is preliminarily added, for the cis-1.4 polymerization of 1,3-butadiene. Such soluble cobalt compound includes those soluble in hydrocarbon-series solvents or riquid 1,3-butadiene to be used or uniformly dispersible ones therein, for example cobalt Pulletone complex, such as cobalt (1) acetylacetonate, and cobalt (1) acetylacetonate, cobalt Pulletone complex, ocheat acetylacetonate, cobalt Pulletone complex, ocheat acetylacetonate, cobalt Pulletone complex, such as cobalt cobate, cobalt high eter complex, cobalt salts of organic carboxylic acids with 6 or more carbon atoms, such as cobalt cototate, cobalt high-there det with acond between the dethyl acond complex such as cobalt chloride cyridine complex and cobalt chloride ethyl acond

complex. The amount of such soluble cobalt compound to be used is preferably 0.001 mmol or more, more preferably 0.005 mmol or more per one mole of 1,3-butadiene. The molar ratio (AVCo) of an organic aluminum chloride to such soluble cobalt compound is 10 or more, particularly 50 or more. Still additionally, organic carboxylate saits of nickel, organic complex saits of nickel, organic complex saits of neodymium and organic complex saits of neodymium and organic complex saits of neodymium may also be used other than the soluble cobalt compound.

[0030] The temperature for the cis-14 polymetization is generally within a range of a temperature above 0°C to 10°C, to 10°C, to 10°C collegation 20°C. Polymerization time (mean retention time) is preferably with a range of 10 minutes to 2 hours. The cis-14 polymerization is preferably done so that the polymer concentration after the cis-14 polymerization may be 5 to 28° by the mass. As the polymerization tank, one tank or two or more tanks in conjugation are used. The polymerization is carried out while the solution is mixed together under agistion in the polymerization tank (polymerization appearatus). As the polymerization tank for uses in polymerization and polymerization appearatus. As the polymerization tank for uses in polymerization. In PS-40-2846, can be used.

[0031] For the VCR production in accordance with the invention, known molecular weight adjusters, for example nonconjugated dienes such as cyclocotadiene, allene and methylallene (1,2-butadiene) or cr-olefins such as ethylene, propylene and butene-1 can be used during the cls-14 polymerization. So as to further suppress gel generation during polymerization, known gelation-preventing agents can be used. Additionally, the content of the cls-1,4 structure in the polymerized product is generally 80 % by mass or more, perferably 90 % by mass or more, with ML10 to 50, preferably 10 to 40 and with substantially no content of gel matters.

20 [0032] 1.3-Butadiene is 1.2 polymerized to produce VCR, by adding an organic aluminum compound represented by the general formula AIR, and earbon suitifies, and the solubie coated compound if necessary as the 1.2 polymerization mixture thus obtained. Then, the resulting 1.3-butadiene may be added to the polymerization mixture obtained. Then, the resulting 1.3-butadiene may be added to the polymerization mixture obtained. Then, the resulting 1.3-butadiene may be added to the polymerization mixture. Otherwise, the resulting 1.3-butadiene may not be added to the polymerization mixture. Otherwise, the resulting 1.3-butadiene may not be added to the polymerization mixture obtained to the polymerization mixture. Otherwise, the preferably includes trimentylustimum, tripicohypitalimum, tripicohypi

2 (0033) The temperature for the 1,2 polymerization is generally within a range of 0 to 100°C, preferably 10 to 100°C, compor preferably 20 to 100°C. The yeld of 1,2-polybudatione can be naised uding 1,2 polymerization, by adding one to 50 parts by mass, preferably one to 20 parts by mass of 1,3-budadiene per 100 parts by mass of the cis-1 4 polymerization mixture to the polymerization system for 1,2 polymerization. The polymerization is preferably within a range of 10 minutes to 2 hours. The 1,2 polymerization is preferably carried outs to that the polymer concentration adjusted the 1,2 polymerization in might be 9 to 25 % by mass. As the polymerization tank, one tank or two or more tanks in conjugation are used. The polymerization is capacity out while the polymerization substitute insket opether under eighation in the polymerization tank (cupted make polymerization tank cupted with an agitation until for highly viscous solution, for example the appearatus described in JP-8-40-2645, can be used, because the viscosity of the polymerization solution is increased during the 1,2 polymerization and the polymer acts to be attached.

[0034] For the VCR production in accordance with the invention, the process of producing VCR by the cis-1,4 polymerization and susbequent 1,2 polymerizations and succeimed above includes a step of adding a polymer substance of a low melting point and with at least one unsaturated double bond per repeating unit to the VCR production system. When the unsaturated polymer abustance is added after VCR production, for example during compounding, the advantage of the invention cannot be obtained. The addition of such unsaturated polymer abustance to a production system is preferably done into the polymerization mixture at an appropriate time point from the cis-1,4 polymerization to the 1,2 polymerization, more preferably at the time of the 1,2 polymerization.

[0035] The unsaturated polymer substance preferably is at least one selected from polyisoprene, crystallizable polybutadiene of a melting point less than 170°C, liquid polybutadiene, a polymeric compound containing oxygen bond, and derivatives thereof.

[0036] Polyisoprene includes for example general synthetic polyisoprene (cis-1,4-polyisoprene at a content of the cis structure of 90 % by mass or more, etc.), liquid polyisoprene, and trans-polyisoprene.

[0037] The crystallizable polybutadiene of a melting point less than 170°C is preferably a crystallizable polybutadiene of a melting point of 0°C to 150°C, which includes for example 1,2-polybutadiene and trans-polybutadiene with low melting points.

[0038] The liquid polybutadiene includes for example polybutadiene with a very low molecular weight and with an intrinsic viscosity $[\eta] = 1$ or less.

[0039] The polymeric compound containing oxygen bond is preferably compounds with ether group, epoxy group,

carboxyl group, ester group, hydroxyl group and carboxyl group. Specific compounds thereof include for example phenol resin, nytion resis, not provided the provided provided and resistance capolymer, polyanyl ether, and allyl ether copolymer. By adding such polymeric compound containing oxygen bond to a polymerization system, the interface affinity changes between cis-polybutadeine as the marks component of the vinyt-cis-polybutadiene rubber and the 1.2-polybutadeine rubber cytest liber cytest of the 1.2-polybutadeine rubber can effectively be improved. [0404] Additionally, the derivatives thereof include for example isoprene-isobytene copolymer, styrene-isoprene-styrene copolymer, styrene-isoprene-styrene book copolymer, isoprene-styrene resistance in the like, and trylorgonated products of these derivatives of these derivatives.

10 [0041] Among the individual unsaturated polymer substances, preferably, isoprene, styrene isoprene-styrene block copolymer and 1,2-polybutadiene of a melting point of 70°C to 110°C are used. Additionally, the individual unsaturated polymer substances may be used shighy or in mixture of two or more thereof.

[0042] When the unsaturated polymer substances as described above are added, the dispersibility of the 1,2-polybutadiene of a melting point of 170°C or more in the cis-polybutadiene nubber as the matrix component can be prominently improved owing to the compatible effect of the unsaturated polymer substance in the resulting VCR, as described above, so that the characteristics of the resulting VCR are so excellent.

[0043] The amount of the unsaturated polymer substance to be added is within a range of preferably 0.01 to 50 % by mass, none preferably 0.01 to 80 % by mass to the obtained why-dis-op-plotudeline nubber. At any time point of addition, further, agitation is done for preferably 10 minutes to 3 hours, more preferably 10 minutes to 30 minutes after addition. In case of a polymeric compound containing oxygen bond, herein, the amount thereof to be added is within a range of preferably 0.01 to 20 % by mass to the obtained viny/cis-polybuatiene rubber. The method for addition in that case is with no specific limitation. During the city-(14 polymerization or 1.2 polymerization or to produce viny/cis polybuatiene rubber, and/or at the termination of the polymerization of viny-les polybuatiene rubber, and/or at the termination of the polymerization of viny-les polybuatiene rubber, and/or at the termination of the polymerization of viny-les polybuatiene rubber, and/or at the termination of the polymerization of viny-les polybuatiene rubber, and/or in the termination of the polymerization of viny-les polybuatiene rubber, and/or in the termination of viny-les polybuatiene rubber, and/or in the viny-less polybuatiene rubber. The viny-less polybuatiene rubber and/or viny-less polybuatiene rubber.

[0044] In addition to the unsaturated polymer substance, an organic compound containing oxygen bond is preferably added. The organic compound containing oxygen bond preferably includes for example compounds with either group, peroxy group, earboxyl group, seter group, hythory group and cadeonly group, which includes for example acid anhydride, aliphatic actoryl acid and the set of t

time is 10 minutes to 30 minutes.

[0045] After the polymerization reaches a predetermined polymerization ratio, known antioxidents are added according to general methods. Prijical examples of such antioxidants include phenol-series 2,6-6-butyl-p-cress (BITT), prosphorous-series trinony/phenyl phosphite (TNP), sultruseries 4,6-bis(potyhthomethyl-p-cresol, and disunyl-3,3-thod-propionate (FID). The antioxidants may be used singly or in combination of two or more thereof. The antioxidents are added to 0.001 to 5 parts by mass per 100 parts by mass of VCR. Subsequently, a polymerization-terminating agent is added to the polymerization system to terminate the polymerization. The method therefore includes for example known methods per se, such as a method of feeding a polymerization mixture after termination of the polymerization or polymerization and changing a large amount of a polar solvent such as activol such as methanol and ethanol and water in the polymerization mixture or introducing inorgenic acids such as other such as a decided such as other such as calcid cognine icadios such as other surface. The polymerization mixture or introducing inorgenic acids such as the polymerization various. The control of the polymerization mixture or introducing inorgenic acids such as sortion such as foreign and calcid such as sortion acid and petucosis calcid, and phytogen choixide gas to the polymerization mixture. Then, the generaled VCR is separated and recovered, rinced and dried according to general methods, to obtain the intended VCR.

[0046] The VCR of the Invention thus obtained generally is at a ratio of the individual components, namely the ratio of 12-polyburdation and melling point of 170°C or more, cis-polyburdatione rubber randite unsaturated polymer substance as described above, where 80 % by mass of the microstructure of des-polyburdatione rubber is cis-14-polyburdatione and the remaining thereof is trans-14-polyburdatione and viryl-12-polyburdatione. The cis-polyburdatione and the unsaturated polymer substance are singly (unsamely, at unreactive states) souble in boiling n-bearing, and the 12-polyburdatione of a melting point of 170°C or more and the unsaturated polymer substance physicallyldhemically adsorbed are insoluble in boiling n-bearing the same place. The remaining has a melting point of 170°C or more and the unsaturated polymer substance physicallyldhemically adsorbed are insoluble in bioling n-bearing city of 170°C to 220°C, and is a crystal fiber in short fiber as described above. Additionally, the ML of cis-polyburdation rubber is 10 to 50, preferably 20 to 40 as described above.

[0047] Additionally, the VCR of the invention is composed of the 1,2-polybutadiene of a melting point of 170°C or more

and the unsaturated polymer substance dispersed uniformly in the matrix of cis-polybutadiene rubber.

[0048] In the VCR of the invention, generally, the 1,2-polybutandene of a melting point of 170°C or more is dispensed in crystal fiber as described above. Additionally, the unsaturated polymer substances can be dispensed in various modes in association with the crystal fiber of the 1,2-polybutadene of a melting point of 170°C or more. As schematically shown in Fig. 1, the various modes include for example a mode of the crystal fiber "2" of the 1,2-polybutadene of a melting point of 170°C or more and the microparticle "3" of the unsaturated polymer substance dispersed in a form being statched to the crystal fiber "2" of the 1,2-polybutadene of a melting point of 170°C or more and the microparticle "3" of the unsaturated polymer substance in a form being statched to the crystal fiber "2" of the 1,2-polybutadene dispersed in a form being attached to the microparticle "3" of the unsaturated polymer substance in the matrix "1" as schematically shown in Fig.3, and a mode of the despersion of the crystal fiber "2" of 1,2-polybutadene in a state thereof included and dispersed in the microparticle "3" of the unsaturated polymer substance in the matrix "1" as schematically shown in Fig.3, and a mode of the despersion of the crystal fiber "2" of 1,2-polybutadene in a state thereof included and dispersed in the microparticle "3" of the unsaturated polymer substance in the matrix "1" is schematically shown in Fig.4, and mode in combination of two or more of the dispersion modes shown in Fig.3, through 4 may be possible. In Figs.1 through 4, "1" expresses matrix, "2", the crystal fiber of the 1,2-polybutadene of a melting point of 170°C or more; and "3", his microparticle of the unsaturated polymer substance in the microparticle "3", his microparticle of the unsaturated polymer substance in the microparticle "3" of the unsaturated polymer substance in the microparticle "3", his microparticle of the unsaturated polymer substance in the microparticle "3" of the unsaturated polymer substance in the microparticle "3" of the unsa

power of 170°C or more; and "3", the microparticle of the unsaturated polymer substance.

[0049] By the method for producing VCRI of the invention, the 1,3-butadiene and the hydrocarbon-series solvent substantially never containing carbon suitide are recovered by separating and removing carbon disulfide from a mother solution of the polymerization microtrecontaining the uneractive 1,3-butadiene, the hydrocarbon-series solvent and carbon disulfide remaining after the separation and recovery of the generated VCR, generally by distillation to separate 1,3-butadiene and the hydrocarbon-series solvent and separation process of carbon disulfide adducts. Additionally, the 1,3-butadiene and the hydrocarbon-series solvent substantially hever containing carbon suifide are recovered by recovering the three components from a mother solution of the polymerization mixture by distillation, and separating and removing carbon disulfide and the hydrocarbon and separation or the separation process of carbon disulfide adducts. The carbon disulfide from the distillate by the adsorption and separation or the separation process of carbon disulfide adducts. The carbon disulfide ror recycling.

[0050] The mass average molecular weight of the boiling n-hexane-soluble matter on a polystyrene basis in the VCR of the invention is preferably 300,000 to 800,000, nor range preferably 300,000 to 800,000, for range or preferably 300,000 to 800,000, for range or preferably 300,000 to 800,000, for range or preferably 100,000 to 800,000,000 to 800,000,000 to 800,000,000 to 800,000 to

period of time, industrially advantageously, at a high caralyst efficiency and with high processability of the catalyst component. Industrially advantageously, at a high conversion ratio, the VCR can be continuously produced, particularly without any attachment to the inner wall or agitation wing inside the polymeration ratio to a part with a side validation (0052). The VCR of the invertion is used singly for use as tire or is used therefor by being compounded with own signation. The synthetic rubber or natural rubber, extended with process oil if necessary and vulcanization with addition of fillers such as carbon black, vulcanizing agents, vulcanization-accelerators and other general compounding agents. With no specific limitation to any tire member, the VCR is used for ided wall, or tread, siftence, bead filler, linner linner, causes, the cord coating, and base tread. With no specific initiation to any tire member, the VCR is used to see the site of the vice of

passite motients.

(DOS3) Compositions prepared by adding the compounding agents to the VCR of the invention and kneading the resulting mixture together have lowered deswell ratios of 20 or less on an index basis, compared with VCR in the related art (lowered such value represents excellency) and have therefore very great extrusion processability.

[0054] Additionally, the VCR composition (compounded material) in accordance with the invention when vulcanized has got improved hardness and tensile stress. The VCR composition has particularly proved 1070-k tensile stress and has the increase around 40 on an index basis compared with the VCR obtained by the methods of the related art (increased such value represents excellency) and a greatly improved reintroling effect. Furthermore, the VCR composition has greatly improved flex-crack-growth around 30 on an index basis (increased such value represents excellency) and exerts an effect of suppressing flex crack. Additionally, the permeability of gases such as oxygen as a thermore interesting the property demanded toward run fatt tie and the like is lowered by around 55 (lower such value represents excellency) on an index basis, compared with the VCR obtained by methods of the related art, exerting an effect on the suppression of heat involved in oxidative deterioration.

[0055] For the exertion of the various physico-chemical properties, preferably, the 1,2-polybutadene crystal fiber dispersed in CNF is partially depresed in a mon-dispersed during more incrofine crystal in the marks of cis-polybutadene crystal fiber unable (abbreviated as "BR" hereinafer) and concurrently present with a large 1,2-polybutadene crystal fiber with an aggregated structure, in other words, the mono-despensed 1,2-polybutadene crystal fiber in the BR matrix is preferably

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in a short fiber of the mean length of the mono-dispersed fiber crystal along short axis being 0.2 µm or less, an aspect ation of 10 or less, the mean number of the mono-dispersed fiber crystal being 10 or more and a meiting point of 170°C or more. In addition to the 1.2 polybutaciene crystal fiber of a meiting point of 170°C or more, preferably, the unsaturated polymer substance preferably, the single finity with the 1.2-polybutaciene crystal fiber in the BR matrix, and is dispersed therein at a state of physical and chemical adsorption in the vicinity of the crystal fiber dispersion of Figs. 2 to 4), as described above, the concurrent dispersion of the 1.2-polybutaciene crystal fiber of a meiting point of 170°C or more and the unsaturated polymer substance in the BR matrix makes the various properties excellent, preferably.

- [0056] A rubber composition prepared by compounding and compounding the VCR of the invention in other synthetic in before or natural rubber is now described in detail. The rubber composition is suitably compounded with 10 is 00 parts, preferably 50 to 200 parts by mass of the VCR per 100 parts by mass of natural rubber, synthetic rubber or a blend nubber at an appropriate ratio thereof. The synthetic rubber preferably includes for example polysoprome rubber and styrene-butadene copolymer tribber. Additionally, a butadiene rubber composition for the can preferably be produced, using the VCR endror a butadiene rubber composition compounded with the VCR.
- 15 [0057] The rubber composition of the invention can be obtained by kneading the individual components using routine Banbury, open roll, kneader and blaxial kneader.
 - [0058] The rubber composition of the invention may be kneaded with compounding agents for routine use in rubber industries, such as vulcanizing agents, valcanization-eccelerators, antioxidants, fillers, process oil, zinc oxide and stearic acid.
- [0059] As the vulcanizing agents, known vulcanizing agents for example sulfur, organic peroxides, resin vulcanizing agents, and metal oxides such as magnesium oxide can be used.
 - [0060] As the valcanization-accelerators, known valcanization-accelerators for example aldehydes, ammonias, amines, quanidines, thioureas, thiazoles, thiurams, dithiocarbamates and xanthates can be used.
- [0061] The antioxidants include for example amine-ketone series, imidazole series, amine series, phenol series, sulfur series and phosphorous series.
 - [0062] The fillers include for example inorganic fillers such as silicic anhydride, calcium carbonate, magnesium carbonate, talc, iron sulfide, iron oxide, bentonite, zinc oxide, diatomaceous earth, china cley, clay, alumina, titanium oxide, silica, and carbon black, and organic fillers such as regenerated rubber and powdery nuber.
- [0063] As the process oil, any of aromatic series, naphthene series and paraffin series may be used.

Examples

compounded as shown below and in Table 1.

[0064] Examples in accordance with the invention are now described specifically below.

35 Example 1

[0065] A solution of 1.6 kg of 1,3-butadiene dissolved in 18 kg of dehydrated cyclohexane was placed in a 30-L stainless steel-made reaction tank with an agitator after the inside was substituted with nitrogen gas, into which 4 mmol of cobalt octoate, 84 mmol of diethylaluminum chloride and 70 mmol of 1,5-cyclooctadiene were mixed, for aditation at 25°C for 30 minutes for cis polymerization. The resulting polymer had ML of 33 and T-cp of 59, and a microstructure of 1,2 structure at 0.9 % by mass, trans-1,4 structure at 0.9 % by mass and cis-1,4 structure at 98.2 % by mass. After the cis polymerization, an unsaturated polymer substance comprising polyisoprene (IR) (ML = 87; cls-1,4 structure at 98 % by mass) was added to the resulting polymerization solution to 5 % by mass (as the percentage to the resulting vinyl-cispolybutadiene rubber), for agitation at 25°C for one hour. Immediately thereafter, 90 mmol of triethylaluminum and 50 mmol of carbon disulfide were added to the polymerization solution, for agitation at 25°C for another 60 minutes, for 1,2 polymerization. After the completion of the polymerization, the resulting polymerization solution was added to 18 liters of methanol containing 4.6-bis(octylthiomethyl)-o-cresol at 1 % by mass, for depositing and precipitating a rubber-like polymerized material, which was then separated and rinsed with methanol, and dried in vacuum at ambient temperature. The yield of the vinyl-cis-polybutadiene rubber thus obtained was 80 %. Subsequently, the vinyl-cis-polybutadiene rubber was treated in boiling n-hexane, to separate insoluble matters and soluble matters, which were then dried. The polymer as a matter soluble in boiling n-hexane had ML of 31, T-cp of 57, and a T-cp/ML relation of about 1.8, where the micro structure was composed of 1.0 % by mass of vinyl-1,2 structure, 0.9 % by mass of trans-1,4 structure and 98.1 % by mass of cis-1,4 structure. Additionally, the mass average molecular weight on a polystyrene basis was 42 x 104, with [η] of 1.7. The number of the mono-dispersed fiber crystal of 0.2 μm or less along short axis as contained in the vinyl-cispolybutadiene rubber was 100 or more per 400 µm², while the aspect ratio was 10 or less and the melting point was 202°C.

[0066] The VCR rubber thus obtained was subjected to physico-chemical assessment after the VCR rubber was

Assessment items and conditions for carrying out the assessment

Kneading method

5 [0067] Kneading was done according to the following procedures.

[Primary compounding]

[0068] Kneading apparatus: Banbury mixer (volume of 1.7 liters)

Rotation number: 77 rpm

Start temperature: 90°C

Kneading procedures:

Time 0: charging VCR/NR (natural rubber)

Time 0: charging filler

Time 3 min: raising ram for cleaning (15 seconds)

Time 5 min: dump.

[0069] The dumped matter was continuously wound with a 10-inch roll for one minute, for round passing three times and subsequent sheet extrusion. After the compound was cooled for 2 hours or longer, the compound was subjected to secondary compounding according to the following procedures.

[Secondary compounding]

26 [0070] After the completion of the primary compounding, secondary compounding was done according to the following procedures.

Kneading apparatus: 10-inch roll

Roll temperature: 40 to 50°C

Rotation interval: 2 mm 30 Kneading procedures:

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- (1) Time 0: winding dumped matter and charging sulfur and vulcanization-accelerator
- (2) Time 2 min: cutting
- (3) Time 3 min: sheet extrusion after corner scraping and round passing

Time period for vulcanization

Measuring apparatus: JSR curelastometer type 2F

Measuring temperature: 150°C

Measuring time period: vulcanization time periods of $t_{80} \times 2$ and $t_{90} \times 3$

Vulcanization conditions

Vulcanizing apparatus: press vulcanization

Vulcanizing temperature: 150°C

[Assessment of physico-chemical properties of raw rubber]

- (55 [0071] The microstructure was analyzed by infrared absorption spectrometry. Based on the absorption intensity ratio at 740 cm⁻¹ for cis, 967 cm⁻¹ for trans and 910 cm⁻¹ for vinyl, the microstructure was calculated.
 - [0072] The Mooney viscosity (ML114) was measured according to JIS K6300.
- [0073] The viscosity in toluene solution (Top) was measured at 25°C, by dissolving 2.28 g of polymer in 50 ml of toluene and using the standard solution for calibrating viscometer (JIS Z8809) as the standard solution and Canon
- Fenske viscometer No. 400.
 M₁₀₀: tensile stress value when a sample of vulcanized rubber exerted an extension ratio of 100 %, as measured
 - according to JIS K6301. $T_{\rm R}$ tensile strength at break of a sample of vulcanized rubber, as measured according to JIS K6301
- [0074] The melting point of the 1,2-polybutadiene crystal fiber was determined on the basis of the peak point on endothermic curve with differential scanning calorimeter (DSC).

[Physico-chemical properties of compounded material]

[0075] Die swell

Measuring apparatus: apparatus for measuring processability as manufactured by Monsanto (MPT)

[0076] Die shape: circle

L/D: 1, 10 (D = 1.5 mm)
Measuring temperature: 100°C
Shear velocity: 100 sec⁻¹

10 [Physico-chemical properties of vulcanized product]

[0077] Hardness, rebound resilience and tensile strength were measured according to measuring methods defined by JIS-K-6301.

[0078] The tanô of dynamic visco-elasticity was measured under conditions of a temperature of 70°C, a frequency of 10 Hz and a dynamic strain of 2 %, using RSA2 manufactured by Reometrics Far East LTD.

[0079] Exothermic property and PS (permanent strain) were measured under conditions of a strain of 0.175 inch, a load of 55 ponds, 100°C and 25 minutes with Goodnich flexometer according to ASTM D623.

[0080] The compressed permanent strain was measured by compression under conditions of a temperature of 70°C and 22 hours with a compression set tester manufactured by Ueshima Seisakusho Co., Ltd. according to JIS K8301 or ASTM D395.

[0081] As the flex-crack-growth, a number of flexing a sample until the crack of the sample reached a length of 15 mm or more with a flexing machine manufactured by Ueshima Selsakusho Co., Ltd. according to ASTM D813 was measured.

[0082] The gas permeability was measured according to the measuring method defined by JIS K7126.

[0083] The tank of the dynamic visco-elasticity was measured under conditions of a temperature of 70°C, a frequency of 10 Hz and a dynamic strain of 2 % using RSA2 manufactured by Rheometrics Far East LTd.

	Tubic	
	Rubber & chemicals	Compounded amount (phr
Primary compounding	VCR/NR	60/40
	HAF carbon	50
	Process oil	10
	Zinc oxide No. 1	5
	Stearic acid	2
	Antioxidant AS	1
Secondary compounding	Vulcanization-accelarator CZ	1
	Sulfur	1.5
,	170.5	

45 Example 2

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[0084] Vinyl-cis-polybutadiene rubber was obtained in the same manner as in Example 1 except for the use of unsaturated polymer substances (additives) to be added as shown in Table 2.

50 Comparative Examples 1 through 4

[0085] Synthesis and compounding were done in the same manner as in Example 1 except for no addition of unsaturated polymer substance (additive), or except for the addition of unsaturated polymer substance not undiregolymerization but during compounding after VCR nubber synthesis (the amount of unsaturated polymer substance to be added was 10 % by mass of VCR).

[0086] Table 2 shows the raw rubber data of vinyl-cis-polybutacliene rubber compositions. In the table, the number of mono-dispersed fiber crystal was the number per 400 μm^2 as an index while such crystal of a length of 0.2 μ or less along short axis was defined as mono-dispersed SPB fiber crystal.

[0087] The microstructure of highly melting SPB in Comparative Example 1 was at 98.8 % by mass of vinyl-1.2 structure, 0.6 % by mass of trans-1,4 structure, 0.6 % by mass of cls-1,4 structure and a ratio (A/B) between (A) matrix BR as a matter isolable in boiling n-hexane and (B) highly melting SPB as a matter isolable in boiling n-hexane was 88/12. In Comparative Example 1, additionally, rspic of the polymer insoluble in boiling n-hexane was 15. (rspic-spressing the magnitude of the molecular weight of 1,2-polybutadiene crystal fiber measured at a temperature of 135°C, and the solvent used was o-dichiorobenzense (10088). In the table, IR represents IR2200 (polyisoprene manufactured by JSR); 1,2-PB represents RB820 (1,2-polybutadiene manufactured by JSR).

	Crystal fiber diameter along major axis	In polymer substance (μm)	0.1 or less	0.1 or less	None	0.5 or more
	Crystal fiber diameter along major axis	in matrix (μm)	0.2 to 0.5	0.5 to 1	1 or more	1 or more
		H.I (wt%)	22.4	22.3	12.1	12.3
	Mono- dispersed SPB	Shape	Particle	Fiber and particle	Fiber	Fiber
01	Mono- dispersed SPB	Aspect ratio	10 or less	10 or less	20 or more	20 or more
Table 2	Mono- dispersed SPB	Fiber crystal number (crystals /400 µm²)	100 or more	50	4	4
		Polymerization solvent (sp value)	Cyclo-hexane (8.1)	Cyclo-hexane (8.1)	Cyclo-hexane (8.1)	Cyclo-hexane (8.1)
	Polymer substance	Time for addition (amountadded in wt %)	At the time of polymerization (10)	At the time of polymerization (10)		At the time of compounding (10)
	Polymer sucstance	Type (melting point)	Œ	1,2-PB (90°C) (90°C)		Œ
			Example 1	Example 2	Comparative Example 1	Comparative Example 2

Examples 3 through 12 and Comparative Examples 3 through 5

[0089] Vinyl-cis-polybutadiene rubber was obtained in the same manner as in Example 1, except for the addition of polymer substances and solvents shown in Table 3.

[0090] In the table, IR represents IR2200 (polyisoprene manufactured by JSR); liquid PB is Hiker CTBN 1300 x 8 (iliquid polybutadiene with a moiecular weight of 3,500 asmanufactured by Ube Industries, Ltd.; yeavylated PB represents Epplead PB3900 (epoxylated polybutadiene with a viscosity of 33 pascal seconds at 45°C as manufactured by Dalcel Chemical Industries, Ltd.; any either polymer is Manyarim AWS-0851 (viscosity of 400 stokes at 100°C as manufactured by NOF corporation).

Table 3

	Polymer	Polymer substance		Mono-dispersed SPB		
	Type (melting point)	Time for addition (amount added in wt %)	solvent (SP value)	Fiber crystal number (crystals /400 μm²)	Aspect ratio	
Example 3	IR	At polymerization (5)	Cyclohexane (8.1)	100 or more	10 or less	
Example 4	Liquid PB	At polymerization (1)	Cyclohexane (8.1)	29	10 or less	
Example 5	Epoxylated PB	At polymerization (0.5)	Cyclohexane (8.1)	100 or more	7	
Example 6	Epoxylated PB	At polymerization (0.5)	Cyclohexane/ benzene = 80/20 (8.3)	60 or more	7	
Example 7	Epoxylated PB	At polymerization (0.5)	Cyclohexane/ benzene = 60/40 (8.5)	30 or more	9	
Example 8	Allyl ether copolymer	At polymerization (1)	Cyclohexane (8.1)	16	8	
Example 9	Epoxylated PB	At Polymerization (1)	n-Hexane (7.2)	20	10	
Example 10	Epoxylated PB	At polymerization (5)	Cyclohexane (8.1)	29	9	
Example 11	Epoxylated PB	At Polymerization (0.5)	Cyclohexane/ n- hexane = 50/50 (7.7)	100 or 10 more	or less	
Example 12	Allyl ether copolymer	At polymerization (0.5)	Cyclohexane/ n- hexane = 50/50 (7.7)	20 or more	8 or less	
Comparative Example 3	-	-	Cyclohexane/ benzene = 80/20 (8.3)	8	11	
Comparative Example 4	-		Cyclohexane (8.1)	4	13	

(continued)

		(
	Polymer	substance	Polymerization solvent (SP value)	Mono-dispersed SPB	
	Type (melting point)	Time for addition (amount added in wt %)		Fiber crystal number (crystals /400 µm²)	Aspect ratio
Comparative Example 5	-	-	Cyclohexane/ n- hexane = 50/50 (7.7)	2	12

[0091] Data of products compounded with vinyl-cis-polybutadiene rubber compositions and vulcanized products thereof are shown below. In Examples 8 through 12 and Comparative Examples 4 and 5, herein, VCR/NI = 1000, representing that NR (natural rubber) was not added during the primary compounding, for compounding.

15 [0092] Smaller indices of 100 sec; gas permeability, exothermic property, PS, compressed permanent strain and tanô show better excellency.
[0093] Larger indices of hardness, M100, TB, EB, TR, Ranborn abrasion, flex-crack-growth and rebound resilience

show better excellency.

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Table 4

			rable	-		
	Example 1	Example 2	Example 3	Example 4	Comparative Example 1	Comparative Example 2
	Physi	co-chemical p	properties of c	ompounded m	aterial (index)	
Die swell	L/D = 1	L/D = 1	L/D = 1	L/D = 1	L/D = 1	L/D = 1
100 sec ⁻¹	70	72	76	85	100	99
	Phy	sico-chemical	properties of	vulcanized pro	xduct (index)	
Hardness	106	107	104	106	100	100
M100	140	139	138	136	100	101
ТВ	107	107	104	107	100	100
EB	102	100	101	100	100	100
TR	103	103	104	103	100	101
Ranborn abrasion (slip ratio: 20 %)	112	109	108	100	100	99
Flex-crack- growth	135	130	136	131	100	104
Gas permeability of N ₂	95	95	95	95	100	100
Gas permeability of O ₂	93	92	93	92	100	100
Rebound resilience	105	104	105	103	100	101
Exothermic properties	87	88	88	89	100	96
PS	82	83	83	84	100	96
Compressed permanent strain	89	88	88	89	100	98
tanδ	86	85	83	84	100	98

Table 5

Example 5 Example 6 Example 7 Comparative Example 3 Physico-chemical properties of compounded material (index) Die swell L/D = 1 L/D = 1 L/D = 1 L/D = 1 100 sec-1 Physico-chemical properties of vulcanized product (index) Hardness M100 ТВ EB TR Ranborn abrasion (slip ratio: 20%) Flex-crack-growth Gas permeability of N2 Gas permeability of O2 Rebound resilience Exothermic properties Compressed permanent strain tanδ

		Table 6		
	Example 8	Example 9	Example 10	Comparative Example 4
Physico-chen	nical propertie	s of compoun	ded material (ii	ndex)
Die swell	L/D = 1	L/D = 1	L/D = 1	L/D = 1
100 sec ¹	73	71	75	100
Physico-che	mical propert	ies of vulcaniz	ed product (inc	iex)
Hardness	107	107	106	100
M100	138	139	140	100
TB	107	107	107	100
EB	102	100	100	100
TR	104	103	103	100
Ranborn abrasion (slip ratio: 20%)	105	106	106	100
Flex-crack-growth	135	129	132	100
Gas permeability of N ₂	95	95	96	100
Gas permeability of O ₂	93	92	94	100
Rebound resilience	103	104	105	100
Exothermic properties	90	91	89	100

(continued)

Physico-chemical properties of vulcanized product (index)						
PS 82 83 82 100						
Compressed permanent strain	86	87	87	100		
Ταπδ	86	83	84	100		

	Table 7		
	Example 11	Example 12	Comparative Example 5
Physico-chemical p	roperties of cor	npounded mate	erial (index)
Die swell	L/D = 1	L/D = 1	L/D = 1
100 sec ⁻¹	70	73	100
Physico-chemical	properties of v	ılcanized produ	uct (index)
Hardness	107	107	100
M100	141	138	100
ТВ	109	107	100
EB	101	102	100
TR	104	104	100
Ranborn abrasion (slip ratio: 20%)	109	111	100
Flex-crack-growth	133	135	100
Gas permeability of N ₂	95	95	100
Gas permeability of O ₂	93	93	100
Rebound resilience	108	107	100
Exothermic properties	86	86	100
PS	79	78	100
Compressed permanent strain	85	87	100
Tanô	80	78	100

[0094] Figs. 5 through 8 are electron micrographs depicting the microstructures of vinyl-cis-polybutadiene rubber actually obtained. Fig.5 is the micrograph of Comparative Example 1, where 1,2-polybutadiene of a melting point of 170°C or more is a whisker-like crystal to form aggregation in the matrix. Fig.6 is the micrograph of Example 3; Fig.7 is the micrograph of Example 2; and Fig. 8 is the micrograph of Example 4, where the aggregation formed by whisker-like crystals in the individual figures is small compared with Fig. 5, indicating better dispersion.

Claims

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- 1. A vinyl-cis-polybutadiene rubber containing 1,2-polybutadiene and a polymer substance of a melting point lower than that of the 1,2-polybutadiene and with at least one unsaturated double bond per repeating unit, where the 1,2polybutadiene and the polymer substance are dispersed at physically and/or chemically adsorbed states in the cispolybutadiene rubber as the matrix component of the vinyl-cis-polybutadiene rubber.
- 2. The vinvl-cis-polybutadiene rubber according to claim 1, where the 1,2-polybutadiene and the polymer substance are dispersed in short crystal fiber and/or particle in the cis-polybutadiene rubber as the matrix component of the vinyl-cis-polybutadiene rubber.

- The vinyl-cis-polybutadiene rubber according to claim 1 or 2, where the 1,2-polybutadiene is 1,2-polybutadiene of a melting point of 170°C or more and the polymer substance is at least one selected from polyisoprene, crystallizable polybutadiene of a melting point of 150°C or less. liquid polybutadiene and derivatives thereon.
- The vinyl-cis-polybutadiene rubber according to any of claims 1 through 3, where the unsaturated polymer substance is contained within a range of 0.01 to 50 % by mass to the total of the crystal fiber of the 1,2-polybutadiene and cispolybutadiene rubber.
- The vinyl-cis-polybutadiene rubber according to any of claims 1 through 4, where the viscosity of the cis-polybutadiene rubber as the matrix component in toluene solution at 25°C is within a range of 10 to 150.
 - The vinyl-cis-polybutadiene rubber according to any of claims 1 through 5, where [n] of the cis-polybutadiene rubber
 as the matrix component is within a range of 1.0 to 5.0.
- The vinyl-cls-polybutadiene rubber according to any of claims 1 through 6, where the content of the 1,4-cis structure
 of the cis-polybutadiene rubber as the matrix component is within a range of 80 % by mass or more.
 - The vinyl-cis-polybutadiene rubber according to any of claims 1 through 7, where the Mooney viscosity of the cispolybutadiene rubber as the matrix component of the vinyl-cis-polybutadiene rubber is within a range of 10 to 50.

- The vinyl-cls-polybutadiene rubber according to any of claims 1 through 8, where the polymer substance is a matter insoluble in boiling n-hexane.
- 10. The vinyl-cls-polybutadiene rubber according to any of claims 1 through 9, where the 1,2-polybutadiene is dispersed in short crystal fiber in the cls-polybutadiene rubber as the matrix component of the vinyl-cls-polybutadiene rubber and the polymer substance is dispersed in particle therein, and where the short crystal fiber of the 1,2-polybutadiene is dispersed in the particle of the polymer substance.
 - 11. The viriy/cis-polybutadiene rubber according to claim 11, where the short crystal fiber of the 1,2-polybutadiene is never contained in the particle of the polymer substance but is also dispersed in the class polybutadiene rubber as the matrix component, and where the length of the short crystal fiber dispersed in the matrix along major axis is within a range of 0.2 to 1,000 µm and the length of the short crystal fiber of the 1.2-polybutadiene dispersed in the particle of the polymer substance along major axis is within a range of 0.0 to 10.5 µm.
- 15. A butadiene rubber composition prepared by compounding a vinyl-cis-polybutadiene rubber according to claim 1 or 2 at 10 to 300 parts by weight her 100 parts by weight and a rubber selected from natural rubber, polyisoprene rubber, styrene-butadiene copolymer rubber, or a blend rubber of at least two Youse thered.
- 13. A butadiene rubber composition for tire, where a vinyl-cis-polybutadiene rubber according to claims 1 through 11, and/or a butadiene rubber composition according to claim 12 is used.
 - 14. A method for producing vinyl-cis-polybutadiene rubber by a step of the cis-14, polymerization of 1,3-butadiene using a cle-1,4 polymerization catalyst in a hydrocarbon-series solvent, a step of the 1,2 polymerization of 1,3-butadiene in the concurrent presence of a 1,2 polymerization catalyst in the resulting polymerization inkture to generate 1,2-polybutadiene of a melting point of 170°C or more, and a step of the separation and recovery of vinyl-cis-polybutadiene rubber generated from the resulting polymerization mixture, the method including a step of adding a polymer substance with at least one unsaturated double bond per repeating unit to the production system of vinyl-cis-polybutadiene rubber.
- 59 15. The method for producing vinyl-cis-polybutadiene rubber according to claim 14, where the polymer substance is at least one selected from polysoprene, crystallizable polybutadiene of a melting point of 0°C to 150°C, liquid polybutadiene, and derivatives thereof.
- 16. The method for producing vinyl-cis-polybutadiene rubber according to claim 14 or 15, where the amount of the polymer substance to be added to the production system is within a range of 0.01 to 50 % by mass to the vinyl-cis-polybutadiene rubber to be obtained.
 - 17. The method for producing vinyl-cis-polybutadiene rubber according to any of claims 14 through 16, where the step

of adding the polymer substance to the production system is carried out in the polymerization mixture at an appropriate time point from the step of the cit. -1, depolymerization step to the step of the separation and recovery of the viryingpolyburadiene rubber generated from the polymerization mixture obtained after the completion of the 1,2 polymerization.

- 18. The method for producing vinyl-cis-polybutadiene rubber according to any of claims 14 through 17, where the hydrocarbon-series solvent is a hydrocarbon-series solvent with a solubility parameter of 9.0 or less.
- 19. A butadiene rubber composition prepared by compounding the viny-les polybutadiene rubber or batined by a production method according to any of claims 14 through 18 at 10 to 300 parts by mass per 100 parts by mass of a rubber selected from natural rubber, polyisoprene rubber, styrene-butadiene copolymer rubber or a blend rubber of at least two types thereof.
- 20. A butadiene rubber composition for tire, where the vinyl-cis-polybutadiene rubber obtained by a production method according to any of claims 14 through 18 and/or a butadiene rubber composition according to claim 12, 13 or 19 is used.

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FIG. 1

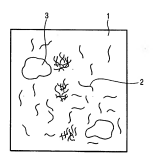


FIG. 2

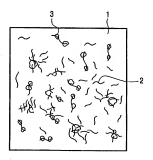


FIG. 3

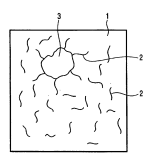


FIG. 4

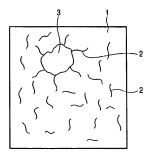


FIG. 5

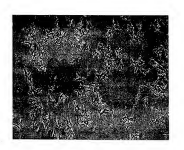


FIG. 6

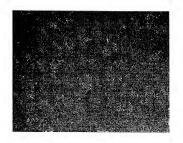


FIG. 7



FIG. 8



	INTERNATIONAL SEARCH REPORT		International applic	sation Mo
	INTERNATIONAL SEARCH REPORT			004/018417
A. CLASSIFIC Int.Cl ⁷	ATION OF SUBJECT MATTER C08L9/00, C08K5/00, C08F2/44,	C08F36/06	101/012	001, 0001
According to Int	ernational Patent Classification (IPC) or to both national	classification and II	·c	
B. FIBLOS SE				
Minimum docum Int.Cl	centation searched (classification system followed by classification syste	ssification symbols) C08F36/06		
Jitsuyo Kokai J	itsuyo Shinan Kcho 1971-2004 Ji	roku Jitsuyo S tsuyo Shinan T	Shinan Koho Toroku Koho	1994-2004 1996-2004
	ase consulted during the international search (name of d	ata base and, where	practicable, search to	rms used)
C. DOCUMEN	ITS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where ap	propriate, of the rele	vant passages	Relevant to claim No.
х	JP 5-194658 A (Ube Industrie: 03 August, 1993 (03.08.93), Claims; Par. Nos. [0001], [00 (Family: none)		0]	1-9,12,13, 19,20
х				1,2,4-9,12, 13,19,20
А	JP B-311246 A (Toyo Tire and ltd.), 26 November, 1996 (26.11.96), Claims (Family: none)	Rubber Co.,	,	1-20
Y Further do	currents are listed in the continuation of Box C.	See patent fi	mily annex.	
"A" document of to be of part "E" earlier applifiling date "L" document to cited to ests special rests "O" document to document to document to the cited to ests."	goined of load documents which is part of the set which is not considered for the part of the set which is not considered foother reference to the part of the set which is stated on present to the published on or after the international which may be three dender on pricely relatingly or which is shifted by the production of the submer denders or other on the specification of the submer denders or other on the specification of the submer denders or other set the submer denders or the submer denders or other means additionally not to the international filling data but have then the relation	"X" document of pe considered no- step when the c "Y" document of pe considered to combined with being obvious	theory underlying the in relicular relevance; the coresid locument is taken alone articular relevance; the commonly on the coresistency.	laimed invention cannot be lered to involve an inventive daimed invention cannot be step when the document is documents, such combination part
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	ng address of the ISAV se Patent Office	Authorized officer		
Facsimile No.	O (second sheet) (January 2004)	Telephone No.		

INTERNATIONAL SEARCH REPORT International application No. PCT/JP2004/018417 C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. JP 2002-338740 A (Ube Industries, Ltd.), 27 November, 2002 (27.11.02), A 1-20 Claims (Family: none) Form PCT/ISA/210 (continuation of second sheet) (January 2004)